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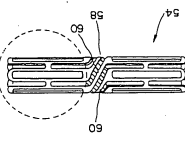
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Stant.

sageway and radial expansion therein is prosageway and radial expansion therein is prowided. The stein (54) is capable of expanding
from a first to a socond diameter by the inelastic
deformation of the material of which the sterl is
comprised. The stent comprises a hoflow tube,
open at both stent ends and having a series of
open at both stent ends on the stent and the
slots (58) therein. The ends of the stent and the
slots (58) therein. The ends of the stent and the
slots are rounded so as to provide smooth
surfaces obviating the abrading of such body **⑤**



EIC.

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25 15 rosclerotic lesion. Once located, the proximal end of blood vessel but instead may be some other body passageway such as the urethra or a bite duct. Curand in particular, to assemblies comprising an outer sheath containing an elongated catheter therein for duct. The assembly is adapted to be percutaneously inserted into a body passageway, somotimes by bly is introduced percutaneously into the femoral artery and then advanced, distally, through the arterial the sheath may be manipulated so as to expose the the intended medical procedure may progress. For example, the so- located distal end of the catheter may include an inflatable balloon for carrying out a procedure. Alternal ively, a prosthesis such as a stent. graft, or stent/graft combination may be delivered, by rently, procedures are performed for stenting such This invention generally relates to assemblies for in a body passageway such as a blood vessel or bite means of a guide catheter. For example, the assemsystem to a desired situs, e.g. at the situs of an athedistal portion of the catheter to the situs, whereafter percutaneous translumenal coronary angioplasty the catheter to such situs. The situs need not be in a delivering devices to a situs in a body passageway delivering the distal portion of the catheter to a situs

and apparatus associated therewith are exemplified by reference to the following U.S. Patents: U.S. Pa-tent Nos. 4,299,226 Issued November 10, 1981 to Banka; 4,323,071 issued April 6, 1982 to Simpson, et al., 4,581,017 issued April 8, 1986 to Sehota; 4,748,982 issued January 7, 1988 to Horzewski, et 81,4,773,899 issued September 27, 1988 to Spears: 4,998,917 issued March 12, 1991 to Gaiser, et al.; 4,998,923 issued March 12, 1991 to Semson, ct el.; 5,007,898 issued April 16, 1991 to Rosenbluth, et al.; 5,034,001 Issued July 23, 1991 to Garrison, et al.; and 4,848,344 issued July 18, 1989 to Sos, et al.: 4,885,003 issued December 5, 1989 to Hillstead; Descriptions of such procedures and the devices 4,932,959 issued June 12, 1990 to Horzewski, et al.; 5,116,309 issued Mary 26, 1992 to Coll.

In carrying out the procedures described and exwhile in some instances, the art has attempted to cure emplified above using heretofore available apparatus, several difficulties have been encountered and, these difficulties, the state of the art is such that fm-

tered is the problem of threading the elongated cathed on the proximal end of the assembly and move the assembly in a distal direction through the passagebly have the requisite stiffness (often termed "pushone is faced with the requirement that the assem-Specifically, one difficulty heretofore

or collapsing. At the same time, the assembly must be led through the tortuous passageway, conforming to This need for both stiffness and conformability is in conflict and such conflict heretofore is manifested in all the bends and turns that are therein encountered. disappointing and unsalistaclory performance of prh

or art devices

Still another difficulty has been encountered in assemblies through the body passageways, there is the employment of the subject devices. In pushing the the great danger of abrading or otherwise traumatidistal direction. Such movement, for example, during cally affecting the inner walls of these passageways The vascular system is particularly vulnerable to such undesirable abrasion. Still further, generally in connection with an emplaced sheath/contained catheter assembly, there is always the danger that the sheath will move relative to the catheter in an undesired direction, such undesired direction being generally the a procedure would obviously be disruptive. Accord-ingly, there is a need to obviate such undesired move-

Summary of the Invention

proved catheters and sheaths are provided which can In accordance with the teachings herein Im-

cooperate to form an assembly obviating the above-described shortcomings of prior devices. In one aspect of this invention a sheath is provideter of such sheath at its proximal portion. Preferably, tion closely adjacent to the distal end and extends for terial employed for such smaller diameter portion is eter, harder polymeric material of construction and larger wall hitchness, is designed to have the requisite "pushability" to transmit forces and translate the ased for containing a device to be delivered to a situs in ployed for the remainder of the sheath. Finally, the ation of smaller diameter, lesser hardness and smallleading distal portion of the assembly as it is being a body passageway e.g., for delivering to such situs the distal portion of a catheter. The sheath comprises an elongated polymeric Lube having an open proximal end and an open distal end and a lumen for contaîning the device, such as a catheter, therein. In accordance with this invention the outside diameter of the sheath at its distal portion is smaller than the outside diamthe smaller diameter distal portion is at only the poronly a small fraction of the length of the sheath at the distal end. Further, the hardness of the polymeric matess than the hardness of the polymeric material emwall thickness of the smaller diameter portion is less than that of the remainder of the sheath. The combinwall thickness results in a flexible, conformable way. On the other hand, the major and lagging proxç 20

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sembly distally through the body passageway. As described herein, all of the above may be accomplished by economically practical manufacturing methods and hence, provides a simple yet highly effective solution to a longstanding problem in this field.

While the differential pushability/conformability of the sheath has been described by a device wherein the diameter, well thickness and hardness of the respective portions have all been varied, it will be understood that a selection of one or more of these parameters may, in certain finstances, produce the desired differential pushability/conformability.

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In another aspect of this invention, an elongated catheter is provided having a proximal and and a distal and. The catheter is adapted to be contained in an elongated tubular sheath for the purpose of having the distal and of such catheter delivered to a situs in a body passageway. The catheter comprises an elongated member having at least one-fumen therethrough, the member having an outer longitudinally extending surface.

in accordance with the teachings herein, the outer surface is provided with a toroidal enfangement in close proximity to the distal end of the catheter. This toroidal enlargement presents, in the fongtudinal cross sectional view of the catheter, a smooth curve, in assembled form, the catheter is contained within the sheath and the linner fumen of the sheath may now be sized such that the distal end of the sheath, in its extrem distal position with respect to the catheth, and have segainst the proximal portion of the toroidal enlargement and hence is precluded from further distal reflocation with respect to the catheter distal realocation with respect to the catheter. Accordingly, the highly undesirable redocation of the sheath during a medical procedure is obviated.

The combination of the new shealh as described above together with the cataleter leadure it is particularly advertageous in that the reduced diameter of the distal portion of the sheath allows such portion to be impaged distally by the endagement without increasing the largest profile of the sheath. That is to say, the endagement may be sized to correspond to the profile of the profile of the produce in the profile of the sheath. That is to the profile of the proximal end of the sheath with the smaller distal end still bearing against the endagement.

In another aspect of this invention, in the specific case of a catheter carrying a prosthesis such as a stent, the same toroidal enlargement placed distally to the stent will prevent the displacement of the stent relative to the catheter.

These and other unique features and benefits of this invention shall be apparent from the following delaifed descriptions and drawings.

rief Description of the Drawings

The invention will be better understood from the following detailed description of exemplary embodi-

ments thereof taken together with the drawing

Figure 1, consisting of Figs. 1A, 18 and 1C, is an elevational, discontinuous view of en assembled sheath and catheter embodying this invention and shown in partlal tongitudinal cross section

Figure 1A is the proximal portion of the assembly including a proximal fitting and a guide wire: Figure 1B is an intermediate portion of the as-

Figure 1B is an intermediate portion of the assembly including an intermediate fitting:
Figure 1C is the distal portion of the assembly, in partial cross section to reveal an inflatable balloon carrying a stent thereupon;

Figure 2 is an enlarged longitudinal cross sectional view of the balloon catheter embodying the teachings of this invention shown in Figure 1C, with the sheath removed and the balloon expand-

ed; Figure 3 is an enlarged, transverse cross sectional view of the portion of the catheler illustral-

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ed in Fig. 2 and taken through line 3-3;
Figure 4 flustrates in longitudinal, elevation view
a stent useful in connection with the assembly IIlustrated in the above drawings; and
Instrated in the above drawings; and
Figure 5 is an enlargement of a portion of the

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Detailed Description of the Invention

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The length of the distal portion 24, in accordance with this invention, is selected to be long enough to conform to the bends and twists of the body passageway. of a device, which in the illustrated embodiment is the tion, is a blood vessel such as a coronary artery. It templated. The sheath 12 comprises an elongated den in Fig. 19) and an open distal end 20, and contains the catheler 16 therein. The sheath is divided Into a relatively long "pushable" proximal portion 22 and a relatively short conformable distal portion 24. example, typically a calheter for carrying a stent to a may range in lengths of from about 35 cm. to about 175 cm. and more typically from about 50 cm. to Referring now to the drawings, illustrated in Figure 1 (Figs. 1Athru 1C) is a sheath/catheter assembly 10 embodying the teachings of this invention. The sheath 12 is designed to deliver the distal portion 14 baltoon catheter 16, to a situs in a body passageway will, of course, be appreciated that other body passageways such as bile ducts or urethras are also conpolymeric tube having an open proximal end 18 (hidthrough which the assembly must be threaded and lead the remainder of the assembly therethrough. For body passageway and passage to the desired situs about 160 cm. The shorter catheters for use in peripheral stenting (e.g., in a femoral or iliac artery) may vary from about 35 cm. to about 90 cm. and the longer which, for the purpose of this specific exemplifica-

catheters for coronary stenting may range from about 90 cm. to about 175 cm. e.g., about 150 cm. The sheath, of course, will be about the same length.

strain, in eccordance with the teachings of this invenlian, it is preferred that the sheath be so divided in distel and proximal portions so that the distal portion is a length of from about 1 cm. to about 35 cm. and more preferably from about 1 cm. to about 12 cm. For example, the distal portion may be 12 cm.

0.005 inches.

As examplified, the distal portion 24 of the sheath 12 kmore condromable then the relatively satisfyeaving a portion, 22 by virtue of having a relatively smaller diameter, a thinner wall thickness and being constructed of a polymer having a lower hardness

will be recognized by those skilled in the art that some stiffness will be required but for all practical purpos-es, a distal portion having the requisite diameter to alremainder of the sheath through the pathway to the desired situs. In contrast with the distal portion 26, the proximal portion 22 is limited in dismeter only by the desire to minimize any trauma to the walls of the low the distal portion of the catheter to slide therein, will have the necessary minimal stiffness to lead the body passageways through which it must pass, except of course, it must retain sufficient flexibility to be passageway trauma will control and practude selecting a diameter for the proximal portion which would be too stiff to manipulate through the pathway. Typiconformability to this leading end of the sheath 12. It portion. Generally, the constraint with respect to body an outside diameter of from about 0.6 mm. (2 French) about 0.5 mm. (2 French) to about 2.3 mm. (7 French). The outside diameter of the proximal portion should The diameter of the distal portion 24 is limited by the distal portion 24 to be easify manipulated to slide over the corresponding distal portion of the devloe. Beyond this limitation, the diameter should be as smail as possible within the practical manufacturing limits so as to present the least trauma and the most lead through the pathway by the conformable distal cally, the distal portion of the sheath may very from to about 6 mm. (18 French) and more preferably. from vary from about 1 mm. (3 French) to about 6.3 mm. (19 French) and more preferably, from about 1 mm. (3 French) to about 2.7 mm. (8 French). For example, the diameter of the distal portion may be 1.55 mm. (4.5 French) and the diameter of the proximal portion the highest profile of the contained device in that it is important that such diameter be large enough to allow may be 1.7 mm. (5 French).

A second contributing factor to the differential pushability/conformability of the disal portion, as congared to the proximal portion, is well thickness; the disal portion having a well thickness less than that of the proximal portion. Sup wall thickness for that of the proximal portion. Sup wall thickness for the disal portion may very from about 0,000 inches and preferably from about 0.0001 inches and preferably from about 0.0001.

inches to about 0.006 inches, for example, 0.003 inches. In contrast thereto, the wall thickness of the proximal portion varies from about 0.006 inches to proximal portion varies from about 0.004 inches to about 0.06 inches and more preferably. from about 0.004 inches to about 0.006 inches, for example,

can be purchased in varying compositions which can result in extruded tubes with varying stiffness. Typh Shore D Duromeler and more preferaby, about 60 to about 70. In contrast thereto, the distal portion is preferably, about 25 to about 60 and more preferably. Stiff a third factor selected for providing the difsheath portions is the hardness of the polymer employed; a hard polymer for the pushable proximal porlion and a soft polymer for the conformable distal porlion. Such polymers as are used currently, generally cally, polymers employed for this purpose are, for example, polyethylenes, polyurethanes, and in some cases, nylons. The polymer of choice is a polyether block polysmide composition sold by the Atochem Corporation of Pennsylvania, under the trade name *PEBAX*. Such PEBAX polymer comes in varying D Durometer values, as the extruded polymer is tested in accordance with the ASTM 1147 standard test mal portion is preferably about 50 to about 70 in hardnesses, ranging from about 25 to about 70 Shore procedure for Shore D Durometer values. The proxiferential pushability/conformability 23 2

As best seen in Fig. 1C, the two portions are joined together by force filling by lea larger diameter portion into the smaller and then "welding" by the application of energy e.g., heat, whereby the polymers fuse to seal the parts alogsther, in can alternative methods to construction the two portions could be continuously co-extruded with polymer of one hardness being first fed to the extruder unit at an upstream station and a polymer of the other hardness being the good to the two portions could be continuously co-extruder with an upstream station and a polymer of the other hardness being fed the downstream thereof.

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about 40 to about 60 in Shore D Durometer value.

body passageway and then the sheath/catheter assembly 10 is threaded over the guide wire 26 by threading such guide wire through a provided guide wire furnen 28, best viewed in Fig. 2. The annular space between the sheath 12 and the catheter is generally flushed with fluid, such as saline solution, to iree the annulus of air which otherwise may be carvia sheath (lush port 39 which is in Intermediate filting 32 and in flow communication with the sheath annulus. The assembly is then advanced through the ster is in the desired position. Referring to Fig. 2. dio opaque markers 30 are provided whereby the pro-Again referring to the drawings, in operation, a wire 26 is generally first introduced into the ned into the body passageway. This is accomplished body passageway until the distal portion of the cathwhich illustrates this distal portion of the catheter, ragress and positioning of the catheter may be monitored by the doctor using x-ray. Once positioned, the guide

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distal portion 22 of the shoath may be drawn back proximally to expose the distal portion of the catheter to the situs. This is accomplished by moving the intermediale fitting 32 proximally relative to the proximal fitting 34. The sheath 12 is affixed to this intermediaffixed to the proximal fitting 34 via a stiffening sec-

ate fitting 32 at its proximal end and the catheter is tion 36. Accordingly, the translation of the intermediare fitting 32 proximally toward the proximal fitting 34 will result in a proximal withdrawal of the distal portion of the sheath from the catheter. This operation is

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of contact with the body passageway as the assembly is being inserted and positioned therein, as contrasted with the blunt end of the sheath, for example. In contrast with the relatively flexible inflatable portion or in the many control is relatively rigid and remains at all itimes in its enlarged configuration. This may be accomplished by manufacturing the balloon integrally with the flange 50 carrying the enlargement by a modding process and varying the flexibility of the life. flatable section from that of the flange by varying the wall thickness of these sections. As Illustrated in the drawings, the wall thickness of the inflatable section of the balcon is shown to be thinner than that of the flange portion. Malerials useful for this purpose are danger of the undesired movement of the distal end dinal cross sectional view shown in Fig. 2, a smooth curved surface. The enlargement is sized relative to within the sheath, the distal end 20 of the sheath, in its extreme distal position as shown in Fig. 1C, bears such undesired movement of the sheath, has the added benefit of providing a smooth non abrading point of the baltoon which must inflate and collapse, the torsuch polymers as ethylene-methacrylic acid polymer, polyurethane, polyethyleneterephthalate, with polyethylene being the material of choice. Alternatively, the enlargement may be made of a similar or dissimilar malerial and attached to the flange by means As described herein, heretofore there has been a Accordingly, the outer surface of the catheter at the flange 50 of the balloon has been provided with a lorthe sheath such that when the catheter is contained against the enlargement 52 and precludes fur ther distal relocation with respect to the catheter. The curved surface of this enlargement, in addition to preduding oidal enlargement 52 which presents, in the longituof the sheath with respect to an emplaced such as gluing, welding or the like. 22

The catheter liself comprises an elongated tube 41 having an outer surface. As exemplified in the drawings and best seen in Fig. 2 and 3, the elongated tube contains a guide wire fumen 28 and a balloon in-

Borst vaive".

end or use elongated tube bych means as werding or gluing. The battoon is made of such a material and is sized such that, in its area of expansion, it is capable of presenting an increased diameter when inflated

by pressure exerted by the introduction of inflating fluid directed to the baltoon via inflation lumen 40

through inflation lumen port 43. When pressure from such inflation fluid in withdrawn, the balloon collaps-

flation luman 40 for carrying fluid to inflate balloon 42. Balloon 42 to circumferentially affixed to the distal

es to a lesser diameter allowing for the retraction of the catheter. Inflation fluid can be introduced into in-flation lumen 40 via inflation fluid port 44 which is

contained within proximal fating 34 and is in flow communication with inflation lumon 40 (see Fig. 14). In the embodiment aboven in Fig. 2, the wire furmen 28 of the alongated tube 41 ferminates at the proximal portion of the balloon. It is necessary for the

guide wire 26 to be threaded through the entire catheter and extend from the distal end thereof as is Illu-

aided by employing the stiffening section 36 in that the catheter lised is generally flexible and manipulation of the catheter is greatly facilitated by such stiffening means. The sheath may be locked fine its position by locking means 36 acrited on the intermediate fitting. Such locking means 38 may, for example, comprise a so-called 'hemostasis valve e.g., a Tuohy

As has been described herein and as is illustrated in Fig. 1C, the balloon may carry an expandable described in U.S. Patent No. 4,733,665 issued March 29, 1988 to Julio C. Palmaz; U.S. Patent No. 4,739,762 issued April 26, 1988 to Julio C. Palmaz; and U.S. Patent No. 5,102,417 issued April 7, 1992 to Julio C. Palmaz and Richard Schalz which are all fincorporated by reference herein. The toroidal enlargement 52, in connection with the placement of such stents is further useful in precluding the undesirable stent 54 for emplacement within a body passageway. Such stents and their delivery and function are well

strated in Fig. 1C. It is also necessary that the entire tumen carrying the guide wire be scaled so as not be in flow communication with the inflating fluid. These goals are accomplished by inserting into the distal

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guide wire In the portion of the catheter extending from the distal portion of the guide wire lumen 28 and

through the distal end of the catheler 48. To Insure flu-

portion of the guide wire fumen 28, a fumen extension 46 which is a fumen containing the

As illustrated in Fig. 4, a stent 54 is provided com-prising a metal tube, open at both ends and having a series of slots 56 therein which allow such stent to be adially expanded into an enlarged diameter by

lumen 28 by adhesive or heat seating means, for example. At the distal end of the balloon, sealing is provided by extending the balloon into a circumferential flange 50 and sealing this flange 50 to the lumen ex-

nside surface of the distal portion of the guide wire

ger of abrading the walls of the body passageway or piercing the balloon of the catheler. Owing to the exshaping of these ends, as well as the shaping of the struts in the articulatable section, is very difficult to defamation of the motal under the action of an expanding force such as the expanding balloon of a balloon catheter. The stent 54 is provided in an interarticulatable section 58. This section links the portion for mability along its length. The stent and its slots are also provided with rounded ends 62, which in contrast to blunt ends, provide a smooth surface with no dantremely small size of the stents (as small as less than mediate position along its longitudinal length with an of the stent with bendable metal struts 60 which can be bent and provide the stent with a degree of conabout 0.5 mm. (less than 2 French) in diameter) the accomplish with conventional equipment. Such shaping may be accomplished by the use of etching techniques but il has been found that the preferred meth-

connection with certain preferred embodiments, it will be apparent to those skilled in the art that various modifications and improvements can be made thereto without departing from the scope thereof. While the invention has been described herein in od is to employ laser cutting apparatus.

 A stent for being radially expanded within a body passageway from a first to a second diameter by the inelastic deformation of the material of which a hollow tube, open at both stent ends and said stent ends and said slots being roundthe stent is comprised; said stent comprising: having a series of slots therein;

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whereby, said stent provides smooth surfaces, obvisting the ebrading of said body passaģ

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The stent of claim 1 wherein said stent is provided with an articulatable section at an intermediate position along the longitudinal length of the stent. ų

The stent of claim 2 wherein the articulatable section comprises bendable struts.

loon capable of providing an expanding force for The stent of claim 1 mounted on an inflatable balsaid radial expansion of said stent.

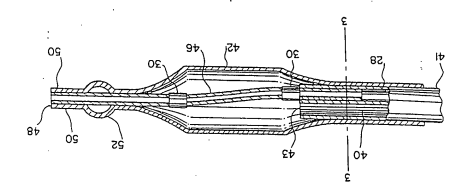
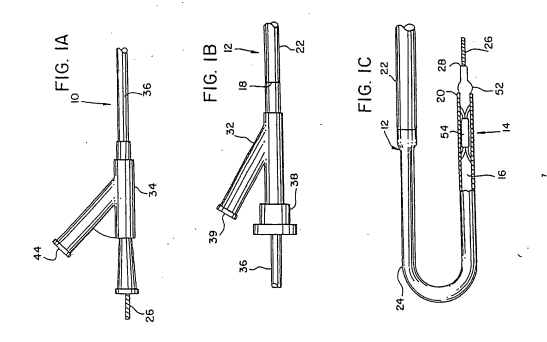


FIG. 2



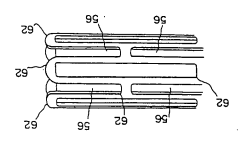


FIG. 5

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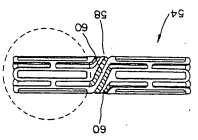
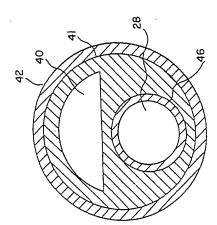


FIG 4



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EUROPEAN SEARCH REPORT

Application Number EP 94 30 0084

	CLASSIFICATION OF THE	A61F2/06				TICGROLL TRUSS SELECTED AG 15.			Mir y Guillén, V
H	Referred to claim	1-4		1-4	1-4			_	Ξ
DOCUMENTS CONSIDERED TO BE RELEVANT	Citation of document with Indiantion, where appropriate, of redering passages	PÁNDÁBLE GRÁFTS	PAUKAZ ET AL.)	VANCED SURGICAL	U ET AL.)		born thems up for all claims	Date of sembidons of the meth	12 April 1994
DOCUMENTS CONS		EP-A-0 335 341 (EXPANDABLE GRAFTS PARTMESHIP) * abstract; floure 7 *	& US-A-5 102 417 (PALMAZ ET AL.)	EP-A-0 274 846 (ADVANCED SURGICAL INTERVENTION) * figures 19,21 *	US-A-5 158 548 (LAU ET AL.) * figures 9-13 *		The present scarch top ort has been strawn up for all claims	The of ores	THE PAGUE
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